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Spread of pathogens through seeds

Occurrence of *Alternaria japonica* on seeds of wild and cultivated rocket

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Abstract

In vitro evaluation were carried out on seed samples of wild and cultivated rocket cultivars most frequently grown in Italy, obtained from farms affected by the leaf spot caused by *Alternaria japonica* in Piedmont and Lombardy during the fall of 2010. Twelve seed samples were collected and assayed for the presence of *A. japonica*. The pathogen was isolated only from not disinfected seeds. Among the two seed samples of cultivated rocket (*Eruca sativa*) only one was infected, at a low level (one infected seed out of 1,600) by *A. japonica*. Four out of ten samples of wild (*Diplotaxis tenuifolia*) rocket seeds were contaminated by *A. japonica* at a frequency of 0.1125 % (nine infected seeds out of 8,000). All isolates of *A. japonica* obtained from seeds were pathogenic on both wild and cultivated rocket.

Key words: seed-borne pathogen; *Diplotaxis tenuifolia*; *Eruca vesicaria*; *Alternaria* black spot

27 **Introduction**

28

29 Wild [*Diplotaxis tenuifolia* (L.) D.C.] and cultivated [*Eruca vesicaria* (L.) Cav.] rocket are
30 popular crops in Italy as well as in many Mediterranean areas, grown for fresh consumption as well
31 as for dish decoration. The two crops have been interested during the past few years by many new
32 soil-borne and foliar diseases, as a result of crop intensification (Gullino et al. 2014). During the
33 fall-winter 2010-2011, extensive necrosis were observed on leaves of *D. tenuifolia* and of *E.*
34 *vesicaria*, grown in commercial plastic-houses in Piedmont and Lombardy (northern Italy). The
35 disease, new for Italy as well as for Europe, is caused by *Alternaria japonica* Yoshii (1941) and
36 interested 30-40% of plants in the affected farms at temperatures ranging between 22 and 25°C and
37 high relative humidity (R.H. >75%)(Garibaldi et al. 2011). The rapid spread of the disease in rocket
38 cultivations suggested that the pathogen could be seed transmitted.

39 The present study was carried out in order to evaluate the contamination of rocket seeds by
40 *Alternaria japonica* in order to consider the possible role of seed transmission in the epidemiology
41 of the disease.

42

43 **Material and methods**

44

45 **Seed infection evaluation**

46 Seed samples of wild and cultivated rocket, belonging to the cultivars most frequently grown in
47 Italy (Table 1), were obtained from farms affected by the disease in Piedmont and Lombardy during
48 the fall of 2010. Twelve samples were collected and assayed for the presence of *Alternaria japonica*
49 on Potato dextrose agar (PDA, Difco, Detroit, Michigan, USA) amended with 25 mg l⁻¹ of
50 streptomycin sulphate as described by Maude and Humpherson-Jones (1980). Subsamples
51 represented by 400 seeds were tested on Petri plates (10 seeds/plate) in two trials. Isolations were
52 made from seeds either non disinfected or surface disinfected for 1 min in 1 % sodium hypochlorite,

53 washed in sterile water for 5 min and dried under a sterile hood. The Petri dishes were incubated at
54 22°C in 12 h light and 12 h darkness at 75% R.H. for 7-10 days. The fungal colonies developing
55 from seeds, morphologically identified as *Alternaria* sp. were transferred from Potato dextrose Agar
56 to Potato Carrot Agar (PCA) and incubated at dark for 7-12 days before the observations at the
57 stereo (Leica,M165C) and optical (NIKON, Eclipse55t) microscopes. *Alternaria japonica* dark-
58 brown colonies on PCA amended with 25 mg l⁻¹ of streptomycin sulphate grew slowly. At the stereo
59 microscope, *A. japonica* colonies presented the fungal mycelium usually grown into the media and
60 conidia produced in singly or in short chains (2-3 elements). Conidia were dark brown, obclavate,
61 obpyriform, ovoid or ellipsoid, with beaks 2-7 (average 3-4) transverse and 1-3 longitudinal septa,
62 and measured 17.7-56.2 (average 30.9) x 6.6-17.8 (average 10.8) µm. Chlamydospores developed
63 usually ten-twelve days after the incubation conditions previously reported. The isolate of *A.*
64 *japonica* from *Brassica chinensis*, coded CBS118390, was used as reference control (Simmons,
65 2007).

66 **Isolates used and their preservation**

67 The isolates obtained from seeds were coded as reported under table 1. Three reference isolates of
68 *A. japonica* were used: AltRuc1/10, obtained from leaves of *E. sativa*, AltRuc 1/11 (Gene Bank
69 accession number JF742643) obtained from infected leaves of *D. tenuifolia* and *A. japonica* coded
70 CBS118390. The different isolates were maintained on PDA at 8 °C.

71 **Inoculum production and pathogenicity test**

72 The different isolates of *Alternaria* spp. were grown on PCA (Potato Carrot Agar) in a growth
73 chamber in darkness at 22-24°C for one week, then kept for another week under 12 hours light /day
74 to stimulate conidia production.

75 For the pathogenicity test, 30-day-old seedlings of cultivated (cv. Coltivata, Franchi) and wild (cv.
76 Grazia, Maraldi) rocket, were transplanted into a steamed potting soil mix (peat: composted

77 broadleaf bark: clay, 60:20:20 vol /vol) in plastic pots (2 L volume) and maintained in growth
78 chamber at 24°C, with 12 hours/day of fluorescent light. Spores and mycelium fragments were
79 removed from the surface of 15-day-old cultures of the different isolates of the pathogen with a
80 spatula. The conidial suspensions obtained were filtered, conidia and mycelial fragments were
81 counted by hemocytometer and adjusted with deionised water to 1×10^6 CFU (colony forming units)
82 ml^{-1} . Five plants of cultivated and wild rocket were inoculated with each isolate by spraying with a
83 suspension of each of the strains obtained from seeds, and covered with plastic bags for 7 days after
84 inoculation. Three reference strains of *A. japonica* were used (Table 2). The pathogenicity test was
85 carried out twice.

86 Not inoculated plants were prepared similarly but sprayed with deionised water. The trials were
87 carried out in a growth chamber at 24 ± 1 °C, under 12 hours /day fluorescent light. Plants were
88 checked two weeks after inoculation for disease development.

89 The data were analysed by univariate ANOVA with Tukey's HSD test using SPSS software 18.0.

90

91 **Results and discussion**

92

93 Among the two seed samples of cultivated rocket (*E. vesicaria*) only one was infected, at a low
94 level (one infected seed out of 800) by *A. japonica* (Table 2).

95 Four out of ten samples of wild (*D. tenuifolia*) rocket seeds were contaminated by *A. japonica*. The
96 pathogen was isolated only from not disinfected seeds (Table 3), at a frequency of 0.1125 % (9 out
97 of 8,000 seeds).

98 From disinfected seeds of cultivated and wilted rocket it was possible to isolate different strains of
99 *Alternaria* spp., but no strain belonging to *A. japonica*.

100 The isolates of *A. japonica* obtained from the different seed lots were coded (Tables 2 and 3),
101 maintained in culture and tested to evaluate their pathogenicity on wild and cultivated rocket.

102 All the plants of wild and cultivated rocket used, belonging to the cvs. Grazia and Rucola coltivata
103 showed typical symptoms of black-brown leaf necrosis surrounded by a yellow halo after
104 inoculation with the isolates of *A. japonica* obtained from seeds. The virulence of the isolates
105 obtained from infected seeds was similar to that of the three reference strains. All isolates affected
106 both wild and cultivated rocket (Table 4).

107 Wild and cultivated rocket were affected during the past few years as a consequence of crop
108 intensification, by a number of new diseases, most of which are seed transmitted (Gullino et al.
109 2014). *A. japonica* is a seed-borne pathogen of plants in the Brassicaceae (David 2002;). This paper
110 provides evidence that also *A. japonica* occurs on rocket seeds, which suggests that rocket seeds
111 may be important in disseminating also this pathogen.

112 The pathogen was isolated only by not disinfected seeds, thus suggesting an external contamination
113 of seeds. At the usual sowing rate for this crop of 2 g seeds per m² corresponding to 1000 seeds /m²
114 for cultivated and 10,000 seeds/m² for wild rocket, also a low percent of contaminated seeds are
115 enough to favour the spread of the disease, as already observed with other pathogens. For instance,
116 in the case of *Plectosphaerella cucumerina* a percent of infected seeds as low as 0.15% may be
117 important in disseminating the pathogen in wild rocket crops (Gilardi et al. 2013 b). Seed infections
118 also contributed to the introduction in Italy of the *Fusarium* wilt agents for the first time observed
119 on both the rocket species (Garibaldi et al. 2004).

120 The occurrence of different species of *Alternaria* on seeds and their role in the spread of diseases
121 was reported also in the case of *Ocimum basilicum* (Gilardi et al. 2013 a), *Brassica oleracea*
122 (Humpherson-Jones and Maude, 1982; Kohl et al. 2010), *Cichorium* spp. (Barreto et al.2008), and
123 *Helianthus annuus* (Jackson et al. 1987).

124 The outbreak of *A. japonica* on rocket represents a new, potential threat to rocket production. The
125 fact that the pathogen infects both wild and cultivated rocket and is seed-borne increases its
126 potential to cause severe losses on a crop intensively grown. Quick and reliable diagnostic tools and
127 seed dressing, carried out with chemical, physical or biological means might help preventing field

128 losses, as already observed in the case of other seed-borne *Alternaria* spp. (Vannacci and Harman,
129 1987).

130

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135

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Code	Host	Cultivar	Seed company
PMP	<i>Eruca vesicaria</i>	Rucola coltivata	PMP SEMENTI SAS, Forlì (Cesena) , Italy
LAS	<i>Eruca vesicaria</i>	Rucola coltivata	La Semiorto, Sarno, (Salerno) , Italy
ORT	<i>Diplotaxis tenuifolia</i>	Ortis	Ortis, Cesena (Forlì-Cesena), Italy
36/Q	<i>Diplotaxis tenuifolia</i>	Venere	T&T Vegetable seeds, Chioggia (Venezia) , Italy
37/Q	<i>Diplotaxis tenuifolia</i>	Giove	T&T Vegetable seeds, Chioggia (Venezia) , Italy
38/Q	<i>Diplotaxis tenuifolia</i>	Frastagliata	Mazzocchi, Casalpusterlengo (Lodi) , Italy
39/Q	<i>Diplotaxis tenuifolia</i>	Brenta	Orosem, Azzano, San Paolo (Bergamo) , Italy
40/Q	<i>Diplotaxis tenuifolia</i>	Reset	Maraldi Sementi, Cesena (Forlì-Cesena), Italy
41/Q	<i>Diplotaxis tenuifolia</i>	Extra	Franchi Sementi Sp.a., Grassobbio (Bergamo), Italy
42/Q	<i>Diplotaxis tenuifolia</i>	Standard (rif. 283/CR)	Olter Sementi, Asti, Italy
43/Q	<i>Diplotaxis tenuifolia</i>	Winter (rif. 71/CB)	Orosem, Azzano, San Paolo (Bergamo) , Italy
44/Q	<i>Diplotaxis tenuifolia</i>	Rucola Selvatica	Olter Sementi, Asti, Italy

172 Table 2 Number of *Alternaria japonica* colonies detected on cultivated rocket seed samples tested

173

Seed sample code	Total number of <i>Alternaria japonica</i> isolates on 800 seeds/sample	
	Not-disinfected	disinfected
PMP	1 (PMP19-NL)	0
LAS	0	0
Total	1 out 1,600 seeds	0 out 1,600 seeds

174

175

176

177 Table 3 Number of *Alternaria japonica* colonies detected on wild rocket seed samples tested

178

Seed sample Code	Total number of <i>Alternaria japonica</i> isolates on 800 seeds/sample	
	Not-disinfected	disinfected
ORT	0	0
36/Q	1(36Q-4NL)	0
37/Q	3 (37Q-16NL; 37Q-13NL; 37Q-22NL)	0
38/Q	3(38Q-1NL; 38Q-9NL; 38Q-19NL)	0
39/Q	0	0
40/Q	0	0
41/Q	0	0
42/Q	0	0
43/Q	2(43Q-1NL;43Q2-NL)	0
44/Q	0	0
Total	9 out 8,000	0 out 8,000

179

180

Table 4 Virulence of different isolates of *Alternaria japonica* obtained from infected rocket seeds, expressed as percentage of infected leaves 21 days after artificial inoculation, evaluated under greenhouse conditions at 24°C. The data are the average of two trials

Isolate code		Obtained from	% infected leaves on			
			Cultivated rocket		Wild rocket	
			cv.Coltivata		cv. Grazia	
AltRuc 1/10	Infected leaves of cultivated rocket	57.5	ab ^a	45.0	ab	
AltRuc 1/11	Infected leaves of wild rocket	45.0	ab	72.5	bc	
CBS118390	CBS collection	52.5	ab	35.0	a	
PMP19	Not disinfected seed	30.0	ab	57.5	a-c	
36Q-4NL	Not disinfected seed	60.0	ab	50.0	a-c	
37Q-13NL	Not disinfected seed	70.0	b	47.5	a-c	
37Q-16NL	Not disinfected seed	65.0	ab	42.5	ab	
37Q-22NL	Not disinfected seed	50.0	ab	57.5	a-c	
38Q-1NL	Not disinfected seed	40.0	ab	32.5	a	
38Q-9NL	Not disinfected seed	40.0	ab	82.5	c	
38Q-19NL	Not disinfected seed	27.5	a	55.0	a-c	
43Q-1NL	Not disinfected seed	57.5	ab	32.5	a	
43Q-2NL	Not disinfected seed	57.5	ab	37.5	ab	

190